

The 3.2 Å Resolution Structure of a Receptor:CheA:CheW Signaling Complex Defines Overlapping Binding Sites and Key Residue Interactions within Bacterial Chemosensory Arrays

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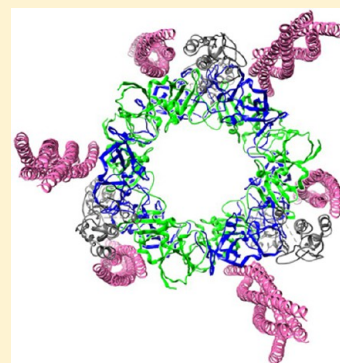
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S Supporting Information

ABSTRACT: Bacterial chemosensory arrays are composed of extended networks of chemoreceptors (also known as methyl-accepting chemotaxis proteins, MCPs), the histidine kinase CheA, and the adaptor protein CheW. Models of these arrays have been developed from cryoelectron microscopy, crystal structures of binary and ternary complexes, NMR spectroscopy, mutational, data and biochemical studies. A new 3.2 Å resolution crystal structure of a *Thermotoga maritima* MCP protein interaction region in complex with the CheA kinase-regulatory module (P4–P5) and adaptor protein CheW provides sufficient detail to define residue contacts at the interfaces formed among the three proteins. As in a previous 4.5 Å resolution structure, CheA-P5 and CheW interact through conserved hydrophobic surfaces at the ends of their β -barrels to form pseudo 6-fold symmetric rings in which the two proteins alternate around the circumference. The interface between P5 subdomain 1 and CheW subdomain 2 was anticipated from previous studies, whereas the related interface between CheW subdomain 1 and P5 subdomain 2 has only been observed in these ring assemblies. The receptor forms an unexpected structure in that the helical hairpin tip of each subunit has “unzipped” into a continuous α -helix; four such helices associate into a bundle, and the tetramers bridge adjacent P5–CheW rings in the lattice through interactions with both P5 and CheW. P5 and CheW each bind a receptor helix with a groove of conserved hydrophobic residues between subdomains 1 and 2. P5 binds the receptor helix N-terminal to the tip region (lower site), whereas CheW binds the same helix with inverted polarity near the bundle end (upper site). Sequence comparisons among different evolutionary classes of chemotaxis proteins show that the binding partners undergo correlated changes at key residue positions that involve the lower site. Such evolutionary analyses argue that both CheW and P5 bind to the receptor tip at overlapping positions. Computational genomics further reveal that two distinct CheW proteins in *Thermotogae* utilize the analogous recognition motifs to couple different receptor classes to the same CheA kinase. Important residues for function previously identified by mutagenesis, chemical modification and biophysical approaches also map to these same interfaces. Thus, although the native CheW–receptor interaction is not observed in the present crystal structure, the bioinformatics and previous data predict key features of this interface. The companion study of the P5–receptor interface in native arrays (accompanying paper Piasta et al. (2013) *Biochemistry*, DOI: 10.1021/bi400385c) shows that, despite the non-native receptor fold in the present crystal structure, the local helix-in-groove contacts of the crystallographic P5–receptor interaction are present in native arrays and are essential for receptor regulation of kinase activity.



Bacterial chemotaxis,¹ the tendency of bacteria to swim toward attractants and away from repellants, has long served as a model system for understanding transmembrane signaling, motility, and cellular behavior.^{2–4} Moreover, the underlying sensory pathways of chemotaxis are essential for the infectivity of many prokaryotic pathogens such as *Helicobacter pylori* (gastric ulcers and stomach cancers),^{5–8} *Vibrio cholerae* (cholera),^{8–10} and several types of pathogenic Spirochetes (lyme diseases, dental disease, syphilis).^{11–13} It has become increasingly apparent that the receptors responsible for binding chemo-

attractants form extended, ordered structures in the cytoplasmic membranes of cells. These chemosensory arrays are primarily composed of the chemoreceptors themselves, also called methyl-accepting chemotaxis proteins (MCPs), the histidine kinase CheA, and the adaptor protein CheW.^{14–18} Cryo-electron microscopy (cryoEM) has revealed a hexagonal arrangement

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for these receptor clusters^{19–24} that is based upon a conserved trimeric assembly of chemoreceptors³ found in species that range from proteobacteria to Thermotogae.²⁵

Although sensing domains differ among MCPs, the receptors all have a similar construction and are exemplified by the four *Escherichia coli* chemoreceptors, Tar, Tsr, Trg, and Tap.^{3,26–28} Dimeric MCPs span the membrane with four helices (TM1, TM2, TM1', and TM2'), bind ligands through a variable amino-terminal extracellular domain, and interact with cellular components through a well-conserved carboxy-terminal cytoplasmic domain (MCP_C). MCP_C is linked to TM2 by a short cytoplasmic HAMP domain that is key to transducing signals across the membrane.^{29–32} Each MCP_C subunit folds as two long antiparallel helices that dimerize into a four-helix bundle.^{33–35} The region most distal to the membrane (the tip of the bundle) known also as the protein interaction region (PIR) or the kinase control module (KCM) interacts with CheA through CheW. In the so-called “adaptation region”, glutamate residues ~140–195 Å away from the receptor tip undergo reversible methylation/demethylation (by CheR and CheB or/and CheD, respectively) to tune receptor activation of CheA.^{36–39} Regulation of the CheB methyl-esterase activity by CheA allows the system sensitivity to adjust to conditions and thereby generates feedback control (also known as adaptation).

The histidine kinase CheA complexes with receptors and transduces ligand binding events into initiation of an intracellular phosphorelay that ends by regulating rotation of the flagellar motor.^{40–42} CheA is a dimer with each subunit containing five separate functional units (P1 to P5), strung together as distinct domains over the length of the polypeptide.^{43–49} P1 contains the substrate histidine autophosphorylated by the kinase domain (P4). P2 docks CheY for phosphotransfer from P1. The last three domains (P3–P4–P5) comprise dimerization, kinase (ATP binding), and receptor-coupling modules, respectively, and their structures have been determined together for *Thermotoga maritima* CheA (CheAΔ289).⁴³

The final core component of the signaling ternary complex is the adaptor protein CheW.^{50,51} CheW has the same tandem SH3-domain-like fold as the CheA P5 regulatory domain and conserves two intertwined five-stranded β-barrels (designated subdomains 1 and 2).^{43,52} The P3-proximal barrel of P5 (subdomain 1) binds CheW through a pseudosymmetric contact that involves conserved hydrophobic residues on each domain.^{34,52–56} Early observations showed that CheW is required for kinase activation,^{57,58} and more recent studies suggest that CheW and CheA P5 may compete for binding receptors.^{59,60}

We produced a model of the receptor:CheA:CheW cytoplasmic ternary complex by application of site-specific spin labeling with nitroxides and pulsed-dipolar electron spin resonance (ESR) spectroscopy to soluble complexes of MCP_C, CheA and CheW from *T. maritima*.⁶¹ Overall, the data revealed that the receptor tip binds CheW but also interacts between the P4 and P5 domains of CheA. The ESR structure is surprisingly asymmetric with the receptor stalk aligning along the CheA dimerization domain and the P1 substrate and P4 kinase domains projected away from the receptor tips,⁶¹ as proposed by a model based on chemical modification studies of the complex.⁵⁴

More recently, we determined the crystallographic structure of a complex between CheA (P4–P5):CheW and a truncated MCP_C from the *T. maritima* soluble receptor Tm14 (PDB code 3UR1⁶²). This structure was fit to electron density from cryoEM maps of native receptor arrays in the cytoplasmic membranes of

cells.⁶² The original crystals diffracted to only 4.5 Å resolution, but their high solvent content enabled placement of the proteins and domains. The receptor PIR interacts with the surface of CheW expected from our PDS studies,⁶¹ but CheA P5 and CheW form rings of pseudohexagonal symmetry that are consistent with the honeycomb receptor lattice observed by cryoEM and predicted by the domain arrangements found by PDS.^{61,62} The combined methods describe a P6 lattice symmetry for the receptor arrays where rings of CheA and CheW associate receptor trimers-of-dimers into a hexagonal network and the P4 kinase domains suspend below the CheW-P5 rings. We have modeled and refined crystallographic structures against the cryoEM data and confirmed that the crystallographic assembly states are consistent with the native arrays within the resolution range of 20–30 Å.⁶² NMR studies verified interfaces implied by the extended structure,^{63,64} and a very similar model has been subsequently published based on independent cryoEM data.²⁴

Despite these advances, the low resolution of the ternary complex structure prevents a detailed description of the molecular interactions within the chemosensory arrays. Indeed, electron density for side chains could not be resolved in the maps, and although the topology of CheA and CheW allow for a largely unambiguous placing of their secondary structure, the register and rotational orientation of the receptor helices was uncertain.⁶² Furthermore, the interaction between the receptor tip and the CheA-P5 domain could only be modeled on the CheW receptor interaction because in the crystals, P5 interacts with the receptor at a position that was only assumed to mimic the native association. Greater structural detail is necessary to not only refine the overall architecture of the arrays, but also understand the mechanism for switching activity states. With the aim of improving the ternary complex crystals, we have reengineered the receptor fragments by perturbing helix termini involved in lattice contacts. One of these altered fragments consistently produced crystals that diffracted to better than 3.5 Å resolution. The resulting higher resolution structure confirms the P5:CheW rings of the previous structure, but the receptor itself displays an unusual unzipped conformation in which a tetrameric 4-helix bundle associates the CheA P5 domains and CheW. This unzipped structure has not been observed in native arrays and is believed to be a non-native feature. However, evolutionary analysis of sequence conservation and mutation patterns suggests that the contacts among the receptor, CheA, and CheW displayed by the structure are relevant to the native chemosensory system. The companion paper directly confirms the predicted receptor-P5 interface is present in the native, membrane-bound array and shows the intact interface is essential for receptor regulation of CheA activity.⁶⁵

■ EXPERIMENTAL PROCEDURES

Constructs of *T. maritima* Tm14_s with altered termini (residues 107–192, 107–193, 107–194, 106–191, 106–192) compared to those that produced the previous 3UR1 structure (107–191) were PCR cloned into vector pET28a (Novagen) and expressed with an N-terminal histidine₆ tag in *E. coli* strain BL21 (RIL DE3) (Novagen) after induction with IPTG at 18 °C and overnight growth for 21 h. The Tm14_s fragments were purified first with Ni-NTA chromatography, followed by overnight thrombin digestion, and then size-exclusion chromatography (Superdex 75 Hi-load FPLC column in 50 mM NaCl, 100 mM Tris pH 7.5, 10% glycerol). *T. maritima* CheW and CheA Δ354 (P4P5 domain, residues 355–671) were expressed and purified as described previously.³⁴

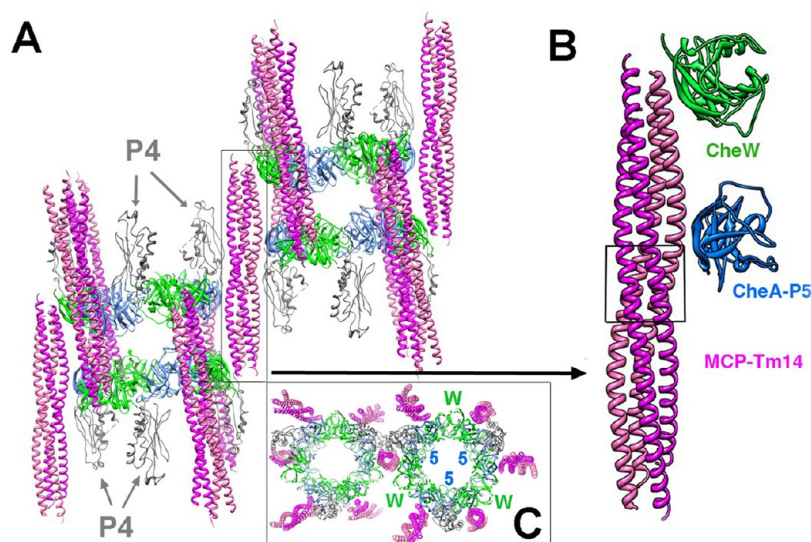


Figure 1. Molecular composition of the 3.2 Å resolution ternary complex crystals. (A) Molecular interactions generated by the R32 crystal symmetry. CheW (green) and the CheA P5 domain (blue) form rings of 3-fold symmetry, with the two proteins alternating around the circumference. The P5-CheW rings are held together by extended 4-helix bundles formed by the Tm14_s receptor (pink and purple). Four Tm14_s subunits have unzipped into continuous helices and associate into a tetrameric 4 helix bundle; each bundle binds two P5 domains near its center (lower site) and two CheW domains at its periphery (upper site). If the pink helices were joined at the middle of the bundle, and the purple helices were similarly joined, they would produce end-to-end MCP hairpin tips as reported in the previous 4.5 Å resolution structure (see Figure 2). Tm14_s tetramers interact with every P5 or CheW domain around the ring, alternating up and down. The CheA P4 domains (gray) project in large solvent channels above and below the rings, although their electron density is not well-defined. (B) Close-up of the Tm14_s tetramer and its interaction with one P5 domain near its center and one CheW at its end. The region that would normally form a helical hairpin instead folds as a continuous helix (boxed). (C) View down the central receptor bundle.

Cubic-shaped crystals ($50 \times 50 \times 50 \mu\text{m}^3$) were grown from a mixture of 520 μM Tm14_s, 457 μM CheA $\Delta 354$, and 121 μM CheW after 1 month by vapor diffusion from a 2 μL drop (1:1 mixture of protein and reservoir: 500 μL reservoir of 0.2 M sodium acetate trihydrate, 0.1 M Tris (pH 8.5), 15% w/v polyethylene glycol 4000). Crystals with a similar shape and size as those derived from Tm14_s (residues 107–191) were grown after 1 month. The new crystals (from Tm14_s residues 107–192) consistently diffracted to 3.5 Å resolution. Crystals were soaked briefly in cryoprotectant that consisted of 85/15% (v/v) reservoir solution with glycerol prior to data collection in a N₂ cold stream. Diffraction data were collected at 100 K with synchrotron radiation at beamline A1 at the Cornell High Energy Synchrotron Source (CHESS). Selenomethionine was also incorporated into the Tm14_s (residues 107–192) to aid in efforts to determine the helical registry, but unfortunately, the selenomethionine incorporated protein did not produce crystals.

Crystal Structure Determination and Refinement. Diffraction data were processed with HKL2000.⁶⁶ Initial phases were obtained with molecular replacement in PHASER⁶⁷ employing the 4.5 Å ternary complex structure with Tm14_s (107–191) (PDB 3UR1) as a search model. The Tm14_s subunits were manually unfolded and built into the resulting electron density with XFIT.⁶⁸ The resulting structure was refined with PHENIX.⁶⁹ B-factor sharpening⁷⁰ and composite omit-map calculation in PHENIX approved maps and allowed for proper interpretation of the Tm14_s unzipping.

Bioinformatics Software and Data Sources. Sequences of CheA, CheW, and MCP proteins were retrieved from the MiST2 database⁷¹ using sets of domain definitions that are specific for each protein, as described previously.⁴⁹ MCP sequences were classified using custom hidden Markov models.²⁸ Pairwise sequence alignments were built using BLAST v.2.2.17⁷² with default parameters. Multiple sequence alignments were con-

structed in MAFFT v.6.0⁷³ with its L-INS-I algorithm. The conservation pattern was analyzed in Jalview⁷⁴ using underlying tools. Minimum evolution and maximum-likelihood phylogenetic trees of CheW protein sequences were constructed from the corresponding multiple sequence alignment and analyzed using the MEGA5 package.⁷⁵ Operons were predicted based on intergenic distances.⁷⁶ A protein cutoff scanning technique⁷⁷ with a β -carbon distance of 6 Å was used to prospect contact sites from PDB files of the cocrystal structures. Measurements were carried out using a custom Perl script, which compares the coordinates of every other β -carbon to the query atom in an all-against-all matrix analysis.

RESULTS

Crystal Lattice Engineering. In efforts to improve the diffraction resolution of the original ternary complex crystals (PDB code 3UR1), modifications were made to the termini of the shortened Tm14 receptor (Tm14_s). The 4.5 Å resolution structure indicated that these termini contacted each other on the symmetry axes of the crystal lattice and thereby allowed the receptor dimers to stack “end-to-end” with aligned helices (Figures 1 and 2). Several new constructs were generated with shifted termini to perturb this principal lattice contact (see Experimental Procedures). One of these (residues 107–192) produced crystals that consistently diffracted to ~ 3.5 Å resolution (Table 1). Ultimately, a new 3.2 Å resolution structure (PDB code 4JPB) was determined from these crystals by molecular replacement (MR) with the previous 4.5 Å structure as a probe (Table 1). MR revealed that the general placement of the CheW, P5, and Tm14_s units were quite similar in the two crystals. Indeed, refinement of the original model against the new data gave $R_{\text{factor}}/R_{\text{free}}$ values of 0.228/0.267 to 3.2 Å resolution. However, examination of the higher resolution electron density (Figure S2 of the Supporting Information) revealed that the helix

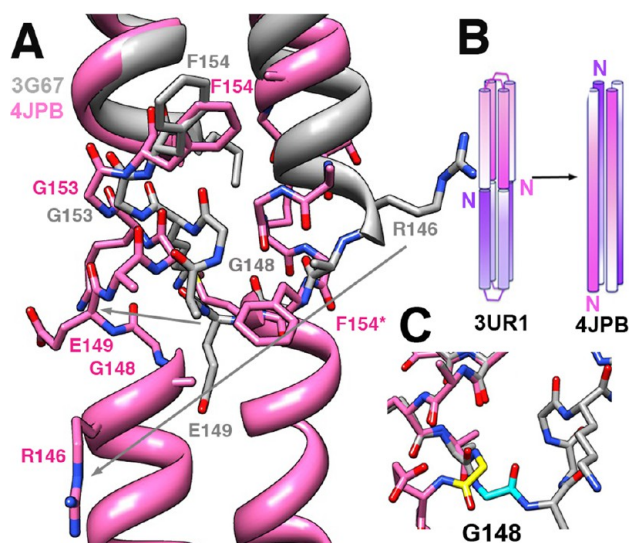


Figure 2. Unzipping of the Tm14_s helical hairpin. (A) Superposition of one subunit of the Tm14 structure (3G67, gray) and two subunits of the unzipped Tm14_s (4JPB, pink). Most of the residues in the transition region maintain similar backbone conformations in both structures, with the exception of Gly 148, whose ϕ/ψ angles change in 3G67 from values disallowed for C β -containing residues to α -helical (see (C)). Gray arrows map residues in the hairpin turn conformation to the extended conformation. The two Tm14_s helices are antiparallel and offset from one another by two helical turns (note the position of Phe154 and its symmetry mate Phe154* in the opposing subunit). (B) Schematic depicting the relationship between two end-to-end hairpins in the previous ternary complex structure of 3UR1 and the tetrameric 4-helix bundle of 4JPB. Note helical hairpins of 3UR1 are inverted relative to that of superimposed 3G67 in (A). Color saturation of the helices decreases from N- to C-termini. (C) Superposition of Tm14 in 3G67 (gray) and in 4JPB (pink) over five residues N-terminal to Gly 148 shows how this residue switches from a Gly-specific conformation (cyan) to α -helical (yellow) in the tetramer.

register in the MR model was not compatible with the side-chain electron density. This was particularly true in the bundle core, where two invariant Phe154 residues appeared to be offset relative to their position in the structure of isolated Tm14 (PDB code 3G67). This consideration led to a new interpretation of the receptor structure, where the subunits had become “unzipped” and then associated into a tetramer of antiparallel helices (Figure 1). Refinement of the new configuration gave significantly improved refinement statistics ($R_{\text{factor}}/R_{\text{free}} = 0.196/0.220$; Table 1). Related fragments of Tm14_s have been studied in complex with CheA and CheW by solution NMR, where they are primarily dimeric.^{63,64} Indeed, Tm14_s appeared largely dimeric on initial size-exclusion chromatography, but analysis of concentrations close to those used for crystallization by multiangle light scattering indicates a mixed species that exchanges between tetramers and dimers (Figure S1 of the Supporting Information). One possible explanation for the unzipping and tetrameric association may be linked to construct design. In addition to the native residues of the receptor, the expression construct introduced four non-native residues at the Tm14_s N-terminus (Gly-Ser-His-Met-Ser₁₀₇). Assuming that Ser107 holds the same position in the helical heptad repeat as it does in the structure of mostly complete Tm14 (PDB code 3G67), the non-native His would reside in a “d” position, internal to the hydrophobic core of the bundle. Two His side chains (one from each subunit) directed at each other in the bundle core

would clash. Thus, their introduction may have destabilized the Tm14_s hairpin dimer under the conditions of crystallization.

Molecular Arrangement within the Ternary Complex Crystal. Despite the switch from a dimeric to a tetrameric Tm14_s, the arrangement of components in this new structure (4JPB) and 3UR1 are quite similar. The conformational changes in the tip that allow for the unzipping of the helical hairpin and the switch from two end-end dimeric hairpins to an antiparallel tetramer of unzipped helices are centered in Gly148 and to lesser extent Gly151 (Figure 2). The Gly148 ϕ/ψ angles change from a region of Ramachandran space disallowed for β -carbon containing residues to the helical region; Gly151 changes conformation more slightly to allow for an i to i+4 main-chain hydrogen bond between Ala150 and Phe154. The Gly rearrangements in the extended tip (residues 147–153) produce a typical heptad repeat of 4-helix coiled-coils with Ala147 and Phe154 residing in the most buried “d” position (Figure 2). Adjacent antiparallel helices of the tetramer pack similarly as in Tm14 or other MCP structures (3G67, 2CH7, 1QU7, 3ZX6), but the symmetry-related helices from the other two subunits are shifted by approximately one helical turn relative to the first pair. This produces a ladder of the four internal Phe154 residues at the center of the bundle. Importantly, the structure presents a pair of antiparallel helices to CheW and P5 as would be found in a typically folded MCP receptor, but the two sides of the bundle (colored purple and pink in Figures 1–5) produce independent binding surfaces offset from one another.

The crystallographic asymmetric unit of 4JPB contains one subunit of CheW and one subunit of the CheA P5 domain, each interacting at a different position on the tetrameric receptor (Figure 1B), with P5 close to the center (lower position; Figure 1B) and CheW at the end (upper position; Figure 1B). Like in the low-resolution structure (3UR1), the R32 crystal symmetry generates rings of alternating CheW and P5 domains (Figure 1AC). Each ring contains three copies of CheW and three copies of P5 (Figure 1C). Each pair interacts through the ends of their β -sheets, with subdomain 1 of P5 binding subdomain 2 of CheW, as previously characterized in complexes of CheA with CheW.^{34,52,62} However, to complete the ring, CheW subdomain 1 also interacts with P5 subdomain 2 in a contact pseudosymmetric to the first (Figure 3). Because CheW and P5 have similar folds, the rings have pseudo 6-fold symmetry. A receptor helix associates with the groove between the two β -barrels of each domain and thereby also produces a pseudo-6-fold symmetric arrangement of receptors. CryoEM in concert with the low resolution structure suggested that in native membrane arrays, one trimer-of-receptor-dimers associates at each P5 or CheW binding site.^{24,62} Because of membrane incorporation of the native receptors, all of the tips would engage the CheW/P5 rings from the same direction, rather than with the alternating polarity found in both 4JPB and 3UR1 (Figure 1). In the unzipped receptor configuration of 4JPB, the Tm14_s helix that primarily binds CheA-P5 does so in a region that would normally be N-terminal to the hairpin tip (this binding helix will henceforth be referred to as the “N-terminal helix”). CheW binds the very N-terminal end of this same helix, but does so with the binding groove flipped over relative to that of P5 (Figure 4). Thus, in the extended crystal lattice, two facing CheW/P5 rings are bound at their edges by six receptor tetramers that alternate their orientation around the rings from “up” to “down” (Figure 1). In 3UR1, weak electron densities for the CheA P4 domains were observed in large pockets below the rings at positions consistent with linkage to P5. In the new structure, the P4

Table 1. Data Collection and Refinement Statistics for Ternary Complex Crystal Structure

wavelength (Å)	0.97700
spacegroup	R32
cell parameters	$a = 213.99$ $b = 213.99$ $c = 208.19$
resolution (Å)	46.2–3.2 (3.3–3.2) ^a
no. of observations	169251
no. of unique reflections	30554
redundancy	5.5 (3.9) ^a
completeness (%)	99.7 (99.2) ^a
R_{merge}^b	0.105 (0.513) ^a
$I/\sigma(I)$	20.3 (1.4) ^a
Refinement statistics	
resolution range (Å)	46.2–3.2 (3.3–3.20) ^a
R factor, %	19.6 (33.6) ^a
R_{free} , %	22.0 (36.9) ^a
molecules/asym unit	1 P4–P5, 1 CheW, 2 Tm14 _s subunits (107–192)
residues/asym unit	576
solvent content (%)	84
overall B-value (Å ²)	36.1
main-chain B-value (Å ²)	32.9
side-chain B-value (Å ²)	39.5
Wilson B-value (Å ²)	40.5
Geometry	
bonds rmsd (Å)	0.01
angles rmsd (°)	1.33
Ramachandran plot, %	
most favored	89.6
additionally allowed	9.5
generously allowed	0.7
disallowed ^c	0.2

^aHighest resolution range for compiling statistics. ^b $R_{\text{merge}} = \sum_i |I_i - \langle I \rangle| / \sum_i I_i$. ^cP5 residue 550 and CheW residue 87 are in disallowed ϕ/ψ regions but have well-defined electron density.

domains localize to the same general positions but appear even less well ordered with their electron densities falling off rapidly beyond the linkage to P5.

CheW Interaction with CheA-P5. In both 4JPB and 3UR1, P5 and CheW produce rings composed of 12 β -barrels, each β -barrel representing a subdomain from either P5 or CheW (Figure 2). At the interfaces formed between subdomains of opposing proteins, three antiparallel β -strands ($\beta 3'$ – $\beta 4'$ – $\beta 5'$ for subdomain 1, and $\beta 3$ – $\beta 4$ – $\beta 5$ for subdomain 2) wrap hydrophobic surfaces against each other in an antiparallel fashion (i.e., the $\beta 3'$ – $\beta 4'$ loop of P5 subdomain 1 interacts with the $\beta 4$ – $\beta 5$ loop of CheW subdomain 2 and $\beta 3'$ – $\beta 4'$ loop of CheW subdomain 1 interacts with the $\beta 4$ – $\beta 5$ loop of P5 subdomain 2; Figure 3). The higher resolution structure (4JPB) shows that Val, Leu, and Ile residues project similarly from $\beta 3'$ – $\beta 4'$ on all four unique surfaces, but the loops connecting the strands differ considerably between the two proteins and also between the two subdomains (Figure 3). The sequence and structural variation within these loops likely gives rise to specificity for ring assembly. In agreement with experiment,^{34,52,78} the interface between P5 subdomain 1 and CheW subdomain 2 is predicted to be stronger (880 Å² buried surface area per subunit; ΔG of formation = –13.6 kcal/mol; specificity $P = 0.024$ ⁷⁹) compared to that between CheW subdomain 1 and P5 subdomain 2 (591 Å² buried surface area per subunit; ΔG of formation = –4.3 kcal/mol; specificity $P = 0.282$). The latter, “weaker” interaction has not been observed outside of the crystallized ternary complexes, although some mutational and modification data suggest that positions on this surface do have a functional role (vide infra).⁵⁵

MCP Interactions with P5 and CheW. The higher resolution structure (4JPB) defines side chain contacts between the receptor helices and P5 or CheW. The junction between the two subdomains of either P5 or CheW harbor branched hydrophobic residues (kinase P5/CheW: kI560/wV27, kI563/wI30, kL547/wL14, kI566/wV33, kL629/wV98) that form a groove to bind the receptor N-terminal helix (Figure 4A,B). The receptor helix binds into this region on P5 with a row of exposed hydrophobic residues (rIle135, rLeu138, rIle142) as well as rAsn139, which provides two key hydrogen bonds to the peptide backbone of kIle566 on the extension of $\beta 2$ that connects the subdomains (417 Å² buried surface area per subunit; ΔG of formation = –5.2 kcal/mol; $P = 0.287$). Mutagenesis studies in *E. coli* have strongly implicated Asn139 in chemoreceptor array structure and function,⁸⁰ and the hydrogen bonding interactions it makes likely serve as an anchor for P5 relative to the receptor tip (Figure 4A). Another potential anchoring contact involves the side-chain to main-chain hydrogen bonds between rArg146 and the kAsp564 at the periphery of the interface. Mutants of the corresponding Arg residue in Tsr (residue 388) were found to impair or abrogate chemotaxis responses in *E. coli*.⁸¹

The interface between CheW and Tm14_s involves the analogous residues on CheW as on P5 (Figure 4B, see above), but the receptor contact forms from a stretch of hydrophobic side chains (rIle109, rLeu113, rIle116) three heptads N-terminal to the P5 contact (331 Å² buried surface area per subunit; ΔG of formation = –4.7 kcal/mol; $P = 0.247$). Moreover, Glu114, which follows the central hydrophobic residue also makes side-chain to main-chain hydrogen bonds with the extension of $\beta 2'$, in

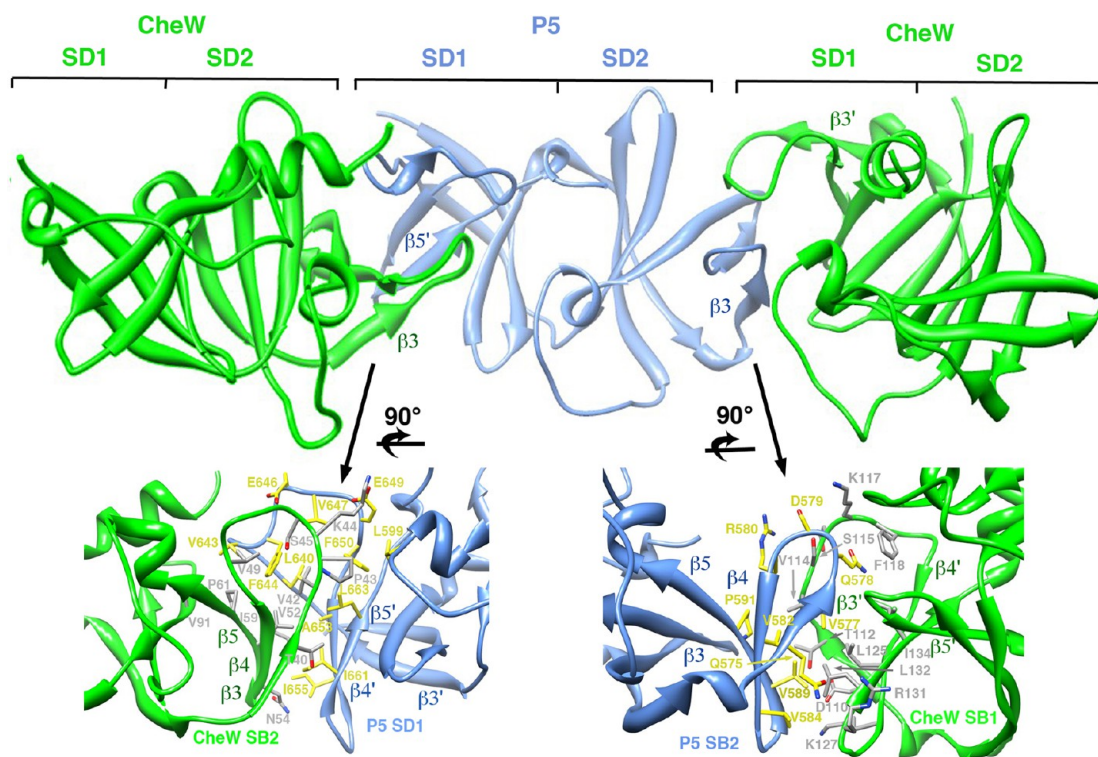


Figure 3. Pseudosymmetric contacts made by the CheW and P5 subdomains. (Top) Half of one P5-CheW ring viewed from the center. (Bottom) The interfaces formed between CheW-subdomain 2 (SB2) and P5-subdomain 1 (SB1) or P5-SB2 and CheW SB1 rotated 90° relative to their orientation above. The two contacts are very similar, with both involving close associations of the $\beta 3$ - $\beta 4$ - $\beta 5$ to $\beta 3'$ - $\beta 4'$ - $\beta 5'$ strands on the respective domains and the conserved hydrophobic residues conserved therein. Nonetheless, substantial differences in the $\beta 3'$ - $\beta 4'$ loops generate specific interactions.

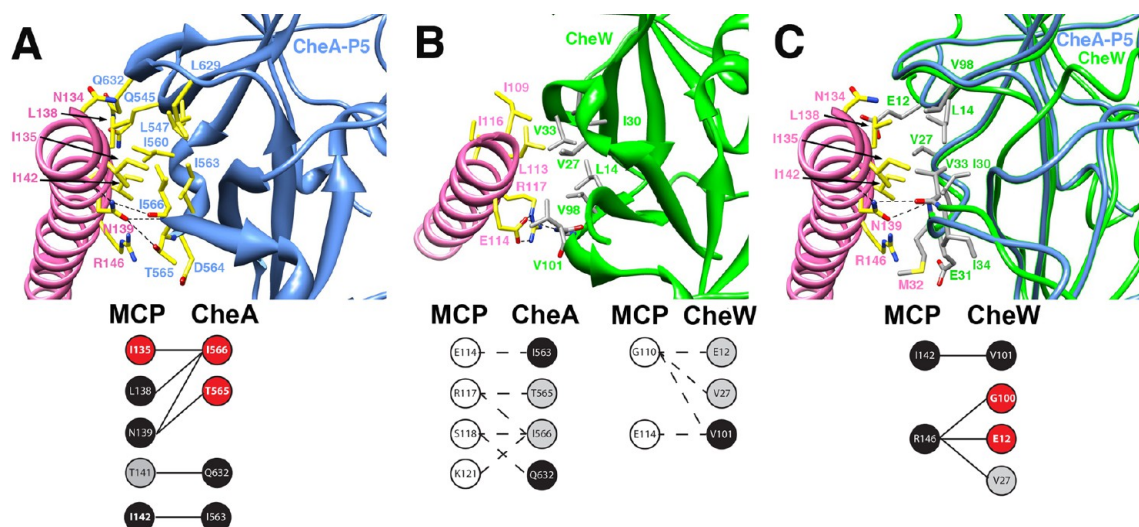


Figure 4. Interactions between Tm14_s and P5 or CheW. (A) Contact between the N-terminal helix of Tm14_s (magenta, N-terminus up) and the groove between subdomain 1 and 2 of CheA P5 (blue). Residues in the interface (yellow bonds) are primarily hydrophobic, with the exception of Tm14_s rAsn139 which hydrogen bonds with the main-chain of P5 kIle566 on the connection between $\beta 2$ and $\beta 3$ and the kThr565 side chain. Below: Binding spots on MCP for CheA predicted by evolutionary information. White circles represent low sequence conservation in a given position (<80% consensus, no functional conservation); gray circles represent strong sequence conservation (>95% consensus, functional conservation); black circles represent the strongest sequence conservation (100% consensus, identical residues or the same charge conservation); red circles represent positions with correlated mutations. Solid lines identify contacts whose evolutionary history is consistent with the correlated mutation hypothesis; dashed lines identify contacts whose evolutionary history is inconsistent with the correlated mutation hypothesis. (B) Contact between the N-terminal helix of Tm14_s and CheW. The Tm14_s helix (pink) runs in the same direction as in (A), but CheW (green) is rotated $\sim 180^\circ$ relative to P5. Because of the pseudosymmetry of the P5/CheW domains, the contact is very similar as in (A), with hydrophobic packing central to the interface and a side-chain-to-main-chain hydrogen bonds between rAsn114, and wVal98, which resides on the connection between $\beta 2'$ and $\beta 3'$. Below: Evolutionary analysis suggests that this receptor site does not constitute a conserved interaction for either CheW or CheA. (C) Superposition of CheW onto P5 demonstrates that CheW conserves chemical character of the residues (gray) at many of the key positions that mediate the contact between Tm14_s and P5. Below: Evolutionary analysis supports this binding mode for CheW. Orientation same as in (A).

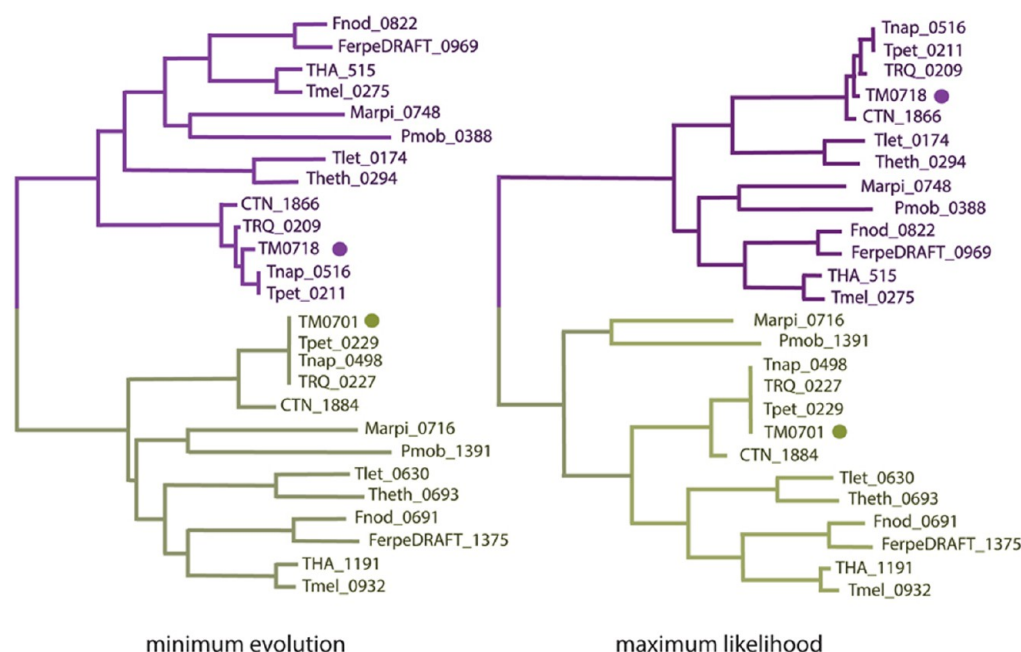


Figure 5. Minimum evolution and maximum likelihood phylogenetic trees showing two groups of CheW orthologs from Thermotogae. CheW1 group is shown in green and CheW2 group is shown in purple. Sequences are represented by their locus tag numbers. Sequences from *T. maritima* are labeled by circles.

close analogy to rAsn139 with $\beta 2$ in the P5 contact. Thus, the upper CheW binding surface of Tm14_s has a similar chemical character compared to the lower surface that engages P5, but the receptor helix runs across the binding groove in the opposite direction. Nonetheless, the intrinsic pseudo-2-fold symmetry of CheW that relates the subdomains essentially compensates for the reversed helix and produces an interaction analogous to that seen with P5 at the lower position.

The upper binding site of CheW, and hence the second P5/CheW ring, is not accounted for by the current models of array structure, which indicate all of the CheW:P5 units are found in one plane, at the tip of the receptors.^{24,62} It is then likely that the upper CheW binding site facilitated crystal lattice formation by mimicking the actual site, which is located at the lower position (residues r135–r146). In support of this notion, rIle135, rIle136, rLeu138, and rIle142 all undergo chemical shifts in solution NMR studies of CheW binding,⁶³ PDS measurements of spin-labeled proteins localize CheW to the lower site,⁶¹ and genetic and biochemical experiments are consistent with this docking arrangement.^{54,82–84} Indeed, superposition of CheW onto P5 indicates an excellent fit with the Tm14_s 135–146 motif into the CheW groove (Figure 4C). However, the strong similarity between the upper and lower receptor binding motifs should not be completely dismissed and raises the possibility of multilayered rings mediated by such contacts in other contexts, perhaps involving receptor systems that are not membrane associated.

Computational Prediction of Protein Binding Sites. We sought to apply a computational genomic approach to independently predict CheW and P5 binding sites on MCP and project what relevance the interactions found in the ternary complex structure may hold in the context of the greater genomic landscape. Rapid accumulation of genomic data in recent years allows productive comparative sequence analyses to identify residues conserved through evolution that are important for structure and function, including protein–protein interactions. A “correlated mutation” hypothesis states that destabilizing

changes in one position can be fixed by a compensatory modification nearby.⁸⁵ The relationship between correlated mutations can be derived from multiple alignments of protein sequences; however, due to the complexity of protein–protein interactions (e.g., mutually dependent residues may not necessarily be in direct contact), there is no single method or approach to successfully predict contact sites. In the case of the chemotaxis system, interacting proteins (MCPs, CheA and CheW) evolve in various subclasses with different protein interaction networks.⁸⁶ This essentially prohibits the application of statistical methods, such as “direct coupling analysis,”⁸⁷ that rely on very large data sets of uniformly interacting proteins. To circumvent this problem and provide direct correlations to the available structural data, we conducted a comparative genomic analysis of the chemotaxis system of *T. maritima* within the well-defined limits of its specific subclass F1⁸⁶ and taxonomic position (phylum Thermotogae).

First, we retrieved sets of CheA, CheW, and MCP protein sequences from all available genomes of organisms from Thermotogae. A comprehensive list of these proteins can be found in Table S1 of the Supporting Information. Satisfactorily, all genomes of Thermotogae contained a single CheA protein that was confidently assigned to the F1 class.

Two Distinct CheW Proteins Are Present in Thermotogae Genomes. We have analyzed the sets of CheW and MCP sequences to reveal potential diversification within these protein families in Thermotogae. Phylogenetic trees constructed from a multiple sequence alignment of CheW protein sequences revealed two distinct sets of orthologs exemplified by *T. maritima* TM0701 (termed CheW1, and the protein contained in the structure presented herein) and TM0718 (termed CheW2): The longest branches on both minimum evolution and maximum likelihood trees separate the two classes (Figure 5). This classification is independently validated by the fact that all *cheW1* genes were found in operons together with *cheA* genes, whereas no *cheW2* genes were a part of these operons. Although this

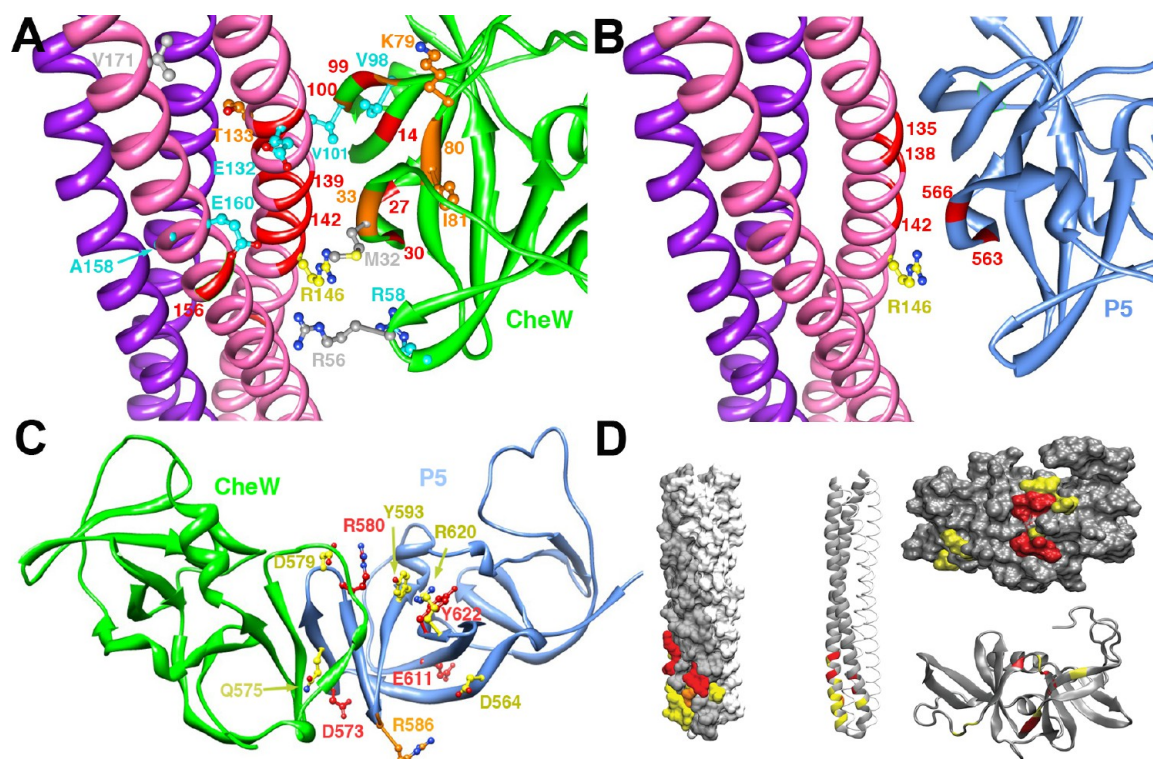


Figure 6. Structure–function analysis of ternary complex interfaces. (A) Residue positions known to report on or affect the interaction between CheW and receptors shown on the model of CheW bound to Tm14_s based on superpositioning CheW onto P5 bound at the receptor tip (Figure 4C). Mutations of CheW residues known to suppress Tsr mutations in *E. coli* are shown as side chains (*T. maritima* numbering). Residue color associates allelic specific suppressors, i.e., mutations at CheW sites that rescue function of mutations at only similar colored sites in the receptor, while at the same time being themselves defective in a wt receptor background (r132 with w98; r133 with w79, w81; r158 with w98, w101; r160 with w57, w58, w101; r171 with w32, w56).⁸⁴ Orange (and red) ribbons represent CheW mutations or modifications defective in receptor interactions (w27, 30, 31, 32, 33, 35, 80, 81, 98, 101).^{82–84,99} Residues identified as mediating Tm14_s-CheW contacts in solution NMR studies shown as red ribbons (w27, 98, 14, 30, 99; r132, 137, 139, 140, 141, 142, 143, 145, 146, 156).⁶³ R146 (yellow bonds) resides at the tip of the MCP hairpin and different residue substitutions at this site can produce locked “on” or “off” kinase behavior. (B) Residues found by solution NMR studies to mediate Tm14–P5 interactions are shown as red ribbons (k563, k566; r135, r138, r142).⁶⁴ (C) Sites of mutation or modification in P5 subdomain 2 that affect function. Residue positions depicted upon mutation or modification produce chemotaxis defects (red side chains; k573, k586, k580, k611, k622), curtail CheW binding (orange side chain r586 and rG587 - not shown) or prevent ligand from deactivating kinase (yellow side chains; k575, k579, k564, k593, k620).^{54–56} (D) Correlation between bioinformatics and NMR data for predicting CheW interaction with MCP. Left, CheW binding sites on MCP (solvent accessible surface and ribbon representations). Right, MCP binding sites on CheW. Residues predicted by “bottom-up bioinformatics” are in red. Residues identified by NMR are in yellow. Overlapping residues are shown in orange.

protein has not yet been experimentally characterized, strong conservation of sequence and structure of the CheW2 protein suggests that this protein is functional in *T. maritima* and all its relatives.

Two Distinct MCP Classes Are Present in Thermotogae Genomes. All 117 MCP sequences from Thermotogae were matched against hidden Markov models (HMMs) constructed for specific MCP signaling classes.²⁸ Eighty sequences were confidently assigned to the 44H (44 helical heptads) class (Table S1 of the Supporting Information), which is the main MCP class within the F1 chemotaxis system.⁸⁶ Further sequence similarity analyses revealed a highly conserved group of 12 MCPs exemplified by the Tm14 protein from *T. maritima* (TM0014). Twenty-four MCPs did not match to any HMM, were sporadically distributed among Thermotogae genomes (indication of horizontal gene transfer) and remain unassigned. The TM0014-type sequences were all composed of a signaling domain of 36 helical heptads, contained no other domains, and showed a very high degree of similarity. On the other hand, in sequence composition they did not match the previously described 36H MCP signaling class;²⁸ therefore, we termed the new class T36H.

One CheW for Each Class of MCPs. We established that genomes of Thermotogae contain two different classes of CheW proteins (CheW1 and CheW2) along with MCPs that also belong to two different classes (T36H and 44H), whereas there was only one CheA protein per genome. On the basis of this finding, we hypothesize that each of the two CheW proteins associates MCPs from each of the two classes with the same CheA protein. If so, which CheW is specific for which MCP class? The structure presented herein, coupled with other substantial experimental evidence, suggests that CheW1 (TM0701) interacts with the T36H MCP (TM0014).^{62,63} Therefore, if our hypothesis is correct, then 44H MCPs should interact with CheW2. There is no experimental evidence to support this claim, because the CheW2 protein from *T. maritima* has been recalcitrant to recombinant production (data not shown). However, we can offer the following computational evidence in support of this hypothesis. It is a well-established fact in evolutionary molecular biology that if two proteins interact they coevolve.⁸⁵ Indeed, CheW1 and T36H proteins appear to be confined to the phylum Thermotogae, whereas CheW2 and 44H proteins are widely distributed throughout the prokaryotes.⁸⁶ For example, the sequence similarity score between CheW1 from

T. maritima and CheW protein from *E. coli* is only 42.4 bits (26% identity), whereas that between CheW2 and the *E. coli* protein is 97.4 bits (37% identity) (Figure S3 of the Supporting Information).

If CheW1 interacts with T36H MCPs and CheW2 interacts with 44H MCPs, then class-specific residues in both protein families (CheW1 versus CheW2 and T36H versus 44H) are candidates for the given interaction. Analysis of the multiple sequence alignments constructed from CheW1 and CheW2 sequences (Figure S4 of the Supporting Information) and T36H and 44H MCPs (Figure S5 of the Supporting Information) revealed several positions in both sets of proteins, where correlated mutations have occurred. For example, a position corresponding to Glu12 in CheW1 (TM0701) is 100% conserved as a negative charge in CheW1 orthologs (in one sequence Glu is changed for Asp); however, the same position in CheW2 orthologs is a positive charge (Lys, 100% conserved). Similarly, the only position, where a reciprocal change is seen in the MCP set is Arg131 in TM0014, which is invariably conserved in all T36H MCPs, whereas all 44H MCPs have a negative charge (Glu, 100% conserved). Thus, according to the correlation mutation hypothesis, corresponding positions in CheW and MCP are mutually dependent; e.g., they may interact. The direct interaction between these two residues is not seen in the ternary complex structure because CheW binds at the upper site in the lattice, most likely due to competition between CheW and P5 for the actual lower binding site. The superposition of CheW onto P5 (Figure 4C) predicts that both residues should reside at the edge of the CheW MCP interface, but will not contact each other. Nonetheless, in functioning arrays CheW and MCPs could sample a range of related association modes, some of which may bring these residues into closer contact. Furthermore, the rArg131–wGlu12 interaction may participate in the recognition of specific receptors by specific adaptors (vide infra). Satisfactorily, this finding is further supported by recent NMR studies, where some of the residues showing significant chemical shift changes upon MCP–CheW binding were identified in very close proximity: rGlu132 (next to rArg131) in Tm14_s and wLeu14 (next to wGlu12) in CheW1.⁶³ Other positions with correlated mutations in CheW (wAsp28, wLys36, wAsp38, wGly100, wLys121) and MCP (rThr141, rAsn159, and rGlu162) are also located in the vicinity of residues that showed significant chemical shift changes upon MCP–CheW binding (Figure 6). Most importantly and in summation, all predicted CheW-binding sites are located at the MCP tip, in agreement with the 3UR1 structure.

“Top-Down” Comparative Genomic Analysis. “Bottom-up” correlated mutation analysis that predicted CheW–MCP interaction sites cannot be used for predicting CheA–MCP interaction sites, because there was only one set of orthologous CheA proteins available for Thermotogae. However, we employed a “top-down” approach, where the evolutionary histories of the interaction sites prospected from the crystal structure were analyzed for consistency with the correlated mutation hypothesis. For example, if a pair of residues in two proteins is predicted to be an interaction site, they should both be well conserved (e.g., invariable, charge preservation, etc) or show a correlated mutation pattern. This approach can be equally applied to MCP–CheA and MCP–CheW interactions, which enabled us to analyze all four putative binding interfaces revealed in two cocrystal structures (Tables S2–S4 of the Supporting Information). Contact sites in interacting proteins were assigned using a protein cutoff scanning technique⁷⁷ and are shown in

Table S2 of the Supporting Information. We then used multiple sequence alignments to trace the evolutionary history of each residue in proposed contacts and analyzed it for consistency with the correlated mutation hypothesis (Tables S3–S5 of the Supporting Information), which states that if contact between two residues is evolutionarily conserved, they will either be invariant or change together in a correlated manner. Summarized results are shown in Figure 4 in reference to the predicted interfaces. Strikingly, in Thermotogae the evolutionary histories of all six MCP–CheA contact pairs and all four MCP–CheW pairs prospected from the top binding site of the MCP are inconsistent with the correlated mutation hypothesis. In a similarly striking contrast, all six MCP–CheA contact pairs and all four MCP–CheW pairs identified at the tip of MCP have evolutionary histories fully consistent with the correlated mutation hypothesis. Furthermore, in both MCP–CheW and MCP–CheA interactions at the tip, there were true correlated mutations. While Arg146 in all MCPs remains fully conserved, its interacting residues in CheW show a correlated mutation pattern. Glu12 and Gly100 are mutually dependent (Figure S4 of the Supporting Information): Glu12 in CheW1 becomes Lys in CheW2, and Gly100 in CheW1 becomes Glu in CheW2. These residues are mutually dependent, most likely because they both interact with the invariable positive charge (residue 146) in MCPs. The pattern of covariance in MCP–CheA interaction is different, but similarly convincing: rIle135 in Tm14 is mutually dependent with kIle566 and kThr565 in CheA. Indeed, rIle135 directly contacts kIle566 in the crystal structure (Figure 4).

Taken together, the evolutionary history of contact sites assigned from the complex crystal structures strongly suggests that (i) both CheW and the P5 domain of CheA bind to the tip of MCP and (ii) MCP contact sites for CheW and CheA binding are not identical, although there is a substantial overlap.

DISCUSSION

The 3.2 Å resolution structure of the *T. maritima* ternary complex differs from the 4.5 Å resolution 3UR1 structure primarily in the modeling of the Tm14_s receptor. First, the 3UR1 structure contains four end-to-end hairpin dimers instead of a tetramer of continuous helices. Second, the location of the CheW/P5 binding surface on the receptor is shifted roughly one turn of helix relative to the current structure. In the higher resolution structure, clear side-chain electron density makes Tm14_s placement certain, whereas in the lower resolution structure, side chains were not discernible, and thus the receptor fragment was placed based on the apparent positions of the termini.⁶² Finally, the polarity of the helical bundles are switched between the structures, which effectively changes the engagement of the lower binding motif from CheW to CheA P5. These considerations raise the question as to whether the lower resolution data would be better modeled by the new structure derived from the higher-resolution data. Difference Fourier maps calculated between the experimental amplitudes from the two data sets show large peaks at the junctions where the hairpins have unzipped, which suggests the structures may be different in this region (Figure S2B of the Supporting Information). Agreement statistics derived from refinement of the old and new models against the low resolution data slightly favor the new receptor configuration, but do not greatly distinguish between the two models ($R_{\text{factor}}/R_{\text{free}}(\text{old}) = 0.207/0.238$; $R_{\text{factor}}/R_{\text{free}}(\text{new}) = 0.206/0.217$). Biochemical analysis of intact arrays, similar to those performed in the companion paper on CheA P5 are needed to verify the detailed CheW Tm14_s interface defined

in 3UR1. Despite this uncertainty, given the arguments presented herein, it is likely that the interactions of receptor with CheW are very similar to those of receptor with CheA-P5. Thus, the overall arrangement of receptors, CheA and CheW in the current array models^{24,62} continues to be supported by nearly all available data.

There is a growing consensus over the structure of membrane associated bacterial chemotaxis receptor arrays, which appear to be universally based on a hexameric assembly of receptors that form trimers-of-receptor dimers, with the greatest ordering at their membrane distal tips where CheA and CheW bind.^{14,22,25,62,88} Recent electron cryotomographic data combined with the modeling of crystallographic structures suggest that CheA and CheW form ring structures of pseudohexagonal symmetry that template the receptors, for at least some states of the arrays.^{24,62} Interdigitated assembly states of overexpressed chemoreceptors, where antiparallel dimers associate through their tips have also been described, but are unlikely to be functional.²¹ The “unzipped” tetrameric assembly for an MCP subunit described here has not been previously observed, and may well result from the shortened Tm14_s fragment that contains several N-terminal non-native residues at its termini. A similar Tm14_s fragment has been used in NMR studies, where the subunits form helical hairpins and behave as typical MCP dimers.^{63,64} Nonetheless, it is worth noting that full-length Tm14 is a naturally “soluble” receptor, in that it has no transmembrane region.³⁵ Computational genomics indicates that CheW1 is specific for this class of receptor (T36H). Soluble MCPs have been observed in other settings, where they have important functions.^{89–92} In some cases these receptors localize to the receptor arrays, whereas in others, they appear to form cytoplasmic clusters.^{89–92} Given the relatively modest conformational changes in the tip that allow for the switch between dimeric and tetrameric states, the possibility that some classes of soluble receptors form extended unzipped structures should not be ruled out. Certainly, substantial rearrangements of coiled-coils play important roles in other systems. Most notably, membrane fusion events involve large-scale conformational changes and repacking of helical proteins. These processes include SNARE mediated vesicle fusion^{93,94} and viral envelope fusion.⁹⁵ In the latter case, viral fusion proteins have hairpin-like helical structures morph into long extended coiled-coil trimers to associate the viral envelope with the plasma membrane.^{95–98} Thus, if soluble MCPs were to unzip and repack, these other processes demonstrate that such changes would not meet a prohibitive thermodynamic barrier.

Despite the unusual tetrameric assembly of Tm14_s, the interfaces found among CheA, CheW, and MCPs in this ternary complex structure are representative of those found in the transmembrane chemosensory arrays. Sequence conservation, as well as the coevolution of interacting sites, strongly supports the groove between subdomain 1 and 2 on both P5 and CheW as being the primary binding location for CheA and CheW on MCPs. For the receptor, the evolutionary analysis only supports the lower, tip-proximal binding site as being the recognition motif for both CheW and P5. Although the current high-resolution structure does not have CheW bound at this position, superposition of CheW with P5 generates an interface that was predicted by computational genomics (Figure 4), is consistent with the hexagonal symmetry of the native receptor arrays, and was anticipated by a large body of additional experimental data. NMR chemical shift perturbations implicate residues on both CheW and Tm14_s that are found within this contact^{63,64} (Figure

6A,D). Pulsed dipolar ESR experiments of spin-labeled ternary complexes of Tm14_s, CheA and CheW combined with targeted disulfide cross-linking also place CheW at the tip of the receptor in close proximity of the lower binding site.⁶¹ In addition, allele-specific mutations of CheW that suppress defective Tsr receptors with mutations near or in the lower binding site map to the CheW binding groove⁸⁴ (Figure 6A). Allele-specific suppressor mutations on two genes can imply that the derived proteins interact with one another. Nonetheless, other coupling mechanisms are also possible. Hence, it is quite remarkable how closely sets of allelic specific Tsr/CheW suppressor mutations localize to the predicted interface between Tm14_s and CheW (Figure 6A). Although mutational studies^{82–84} and chemical modification/protection^{54,99} also support the subdomain 1–2 groove on CheW as being the primary MCP binding site (Figure 6A), the suppressor studies genetically link this recognition motif directly to the corresponding surface on the MCP tip.⁸⁴ It should also be noted that there is one set of CheW/Tsr allelic suppressors that map closest to one another when only the upper binding site on Tm14_s is considered (Figure S7 of the Supporting Information). Although it is tempting to interpret this data as evidence for functional relevance of the upper binding site, it is more likely that longer-range structural coupling through the receptor propagates the effects of this mutation to CheW binding at the lower position.

Several lines of evidence suggest that the lower binding site on Tm14_s of the current structure represents the primary association mode of CheA with receptors. Specific residues on the receptor tip, where substitutions greatly perturb chemotaxis in *E. coli*, play critical roles in the interface with P5 at the lower site. For example, rAsn139 makes key hydrophilic contacts with partner proteins by hydrogen bonding directly with the main chain of kIle566 and wVal33 (Figure 4). Substitutions of the analogous residue in Tsr (Asn381) have dramatic functional effects, with all but a Gly substitution destroying chemotaxis.⁸⁰ In the context of the hexagonal arrays, Asn381 is also predicted to mediate receptor trimerization.⁸⁰ Thus, the near essential nature of this residue results from its participation in three distinct interfaces: those with CheA, CheW and two other receptor subunits. In another case, rArg146, which resides at the base of the lower interface (and at the boundary of the tip; Figure 4), hydrogen bonds with the main-chain of the connection to subdomain 2 and is close to forming a salt-bridge with kAsp564 (wGlu31). Substitutions to large residues at this site in Tsr (Arg388) produce either “lock-on” (Trp, Tyr) or “lock-off” (Phe, His) kinase activity.⁸¹ Because of its potential to also mediate receptor trimer contacts on the adjacent subunit, these phenotypes also likely result from combined effects at multiple interfaces. Nonetheless, the mutational studies do suggest that alterations in structure at the interfaces resolved by the current structure could be critical for controlling kinase activity.

Despite considerable genetic and biochemical studies of CheA, there has been less direct functional data implicating the P5 surface that binds Tm14_s in function.^{54–56} This is probably because mutations or modifications at many sites affect P5 structure and CheW binding, and these properties are coupled to receptor interactions.^{54–56} Nonetheless, NMR studies that rely on methyl-TROSY experiments of *Thermotoga* proteins identified the same interface defined by the structure and predicted by computational genomics (Figure 6B,D). Most importantly, the companion paper to this report⁶⁵ has taken a targeted disulfide cross-linking and mutagenesis (TAM-IDS) approach to define the CheA–receptor contacts in both isolated

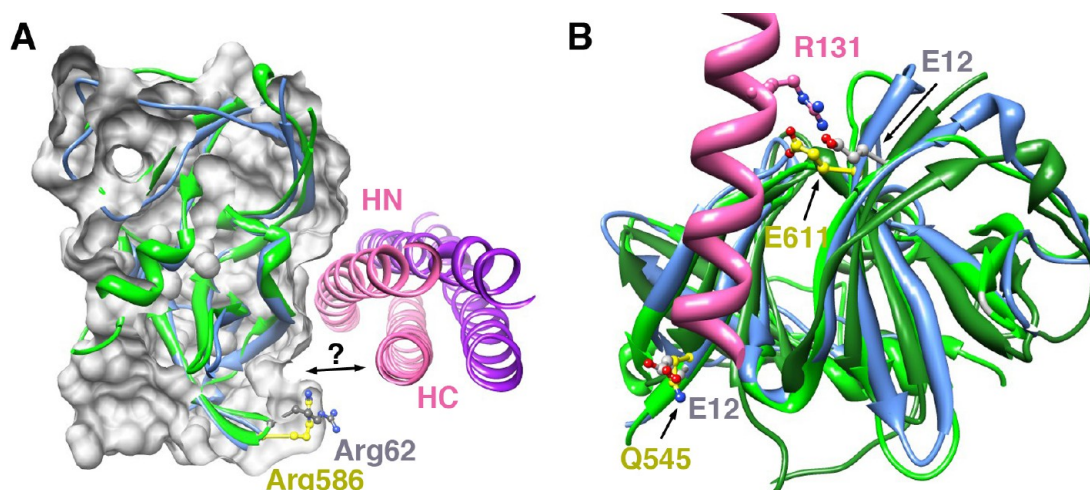


Figure 7. Hypothetical alternate interfaces in chemosensory clusters. (A) wArg62 (kArg586) is a mutational hotspot on CheW, but does not directly engage the receptor, which binds mainly through the N-terminal helix (pink, HN). A relative rotation of CheW (green ribbons, gray side chain), P5 (blue ribbons, yellow side chain) or receptor could engage subdomain 2 with the C-terminal helix (HC) in some states of the ternary complex. (B) wGlu12 (kQ545) and rArg131 show correlated changes in residue identity. If CheW subdomain 1 is superimposed with P5 subdomain 1 (CheW - light green ribbons, gray side chains, P5 - blue ribbons, yellow side chains) to maintain a similar directionality of helix binding, wGlu12 and rArg131 (pink) both participate in the interface, but do not contact. However, if subdomain 1 of CheW is superimposed with P5 subdomain 2 (CheW - dark green ribbons, gray side chains), wGlu12 corresponds to kGlu611 (yellow side chain) and would be positioned to salt-bridge with rArg131.

and cellular chemosensory arrays. These methods were able to distinguish the current interface from the one modeled on the 4.5 Å resolution structure and verify with considerable detail that the interface defined by the higher resolution structure functions in native arrays. Thus, P5 binds at the lower site on receptor tip in a similar orientation to CheW, and this at least partially explains the competition of CheA and CheW for overlapping sites on receptors.^{59,100} The companion paper also shows that the receptor-P5 interaction plays an important role in receptor-mediated regulation of kinase activity.

Both ternary complex crystal structures contain large rings formed by the P5 and CheW subdomains that have been presumed to template receptor trimers in hexagonal arrays.^{24,62} The ring contact between P5 subdomain 1 and CheW subdomain 2 contact has been characterized by a variety of approaches.^{34,52,54–56,99} The secondary contact that completes the ring structures (subdomain 1 of CheW to subdomain 2 of CheA) has not been previously observed outside of crystal structures; however, mutational data and chemical modification experiments have implicated residues near this interface in function (Figure 6C). There are numerous sites in P5 subdomain 2 that when modified produce chemotaxis defects or affect CheW binding; however, many other modifications in subdomain 1 produce similar outcomes.^{55,56} Notably, some Cys-substitutions (and subsequent modification) in subdomain 2 generate phenotypes in which CheA does not deactivate properly with chemoattractant.⁵⁵ Such behavior results only from subdomain 2 substitutions, and several reactive sites localize directly to the interface between subdomain 2 and CheW subdomain 1 (Figure 6C). Thus, modulation of this ring contact may play an important role in kinase regulation.

Despite our advancing understanding of the overall architecture and interactions within chemosensory arrays, there are many details to be resolved. Array function must involve transitions among different structural states that produce different levels of kinase activity. Recent computational work suggests that hexagonal lattice models may correlate with the active state of CheA, although this remains to be verified.¹⁰¹

Given what we know about the assembly modes and the large amount of biochemical and genetic data available on the chemotaxis system, can we infer additional interactions not visualized in the current structures? Although caution must be taken in the interpretation of mutational data due to the networked, potentially redundant nature of molecular interactions within the arrays, a few observations deserve note. For example, substitutions or modifications of wArg62 (*E. coli* CheW Arg56) dramatically affect MCP binding and chemotaxis;⁸³ however, this residue does not directly contact the receptor bundle, despite being oriented toward the C-terminal helix (Figures 4 and 7). Furthermore, mutations of Tsr that suppress CheW mutations at Arg62 and at other residues on the same β 4- β 5 loop that contains Arg62 map to exposed positions on the C-terminal helix of Tm14s⁸⁴ (Figure 6A). Notably, Arg62 is also conserved as an Arg on P5 subdomain 2 (kArg586). Thus, in some states of the array, a relative rotation or deformation of the components may engage the end of subdomain 2 from CheW and/or P5 with the C-terminal helix of the receptor (Figure 7A). Indeed, the companion paper provides the first evidence that such a rotation occurs as targeted disulfide cross-links between kT565 (*S. typhimurium* E550C) and rI156 (Tsr V398C) form more rapidly in the presence of chemoattractant.⁶⁵ This change in reactivity is consistent with a closer association of P5 subdomain 2 and the receptor C-terminal helix when ligand inhibits CheA.

The correlated residue changes at w12 and r131 identified by bioinformatics in the two *T. maritima* CheW/receptor classes also reveal an intriguing structural relationship. The equivalent residue to wGlu12 in P5 does not contain a carboxylate (kGln545), but the pseudo symmetric residue on subdomain 2 does (kGlu611). kGlu611 makes a direct salt bridge with Arg131, which is the position that undergoes a correlated switch to Glu when w12 changes to Lys. Thus, if the lower binding site were to engage CheW with the N-terminal helix running in the opposite direction, w12 and r131 would salt bridge (Figure 7B). Note that this is the orientation that the N-terminal helix takes with respect to CheW at the upper nonconserved site (Figure 4B). At least for

CheW and Tm14 in solution, PDS and targeted disulfide cross-linking favor the CheW orientation that mimics that of P5.⁶¹ Nonetheless, it may be possible in some instances for the CheW/P5 binding groove to accept a receptor helix with either polarity. An inverted arrangement that satisfies the w12-r131 pair would not be compatible with a membrane-associated array, where the receptors all project toward CheA/CheW from the same direction, but such constraints are not present for naturally soluble receptors clusters. Thus, the symmetry of their architectures could be different than those of the membrane arrays and may involve elaborations of the interfaces found in the current structures.

■ ASSOCIATED CONTENT

Supporting Information

Multangle light scattering analysis of Tm14s; electron density maps of ternary structure; pairwise sequence alignments among CheW proteins from various species; sequence alignments of Thermotogae CheW proteins; sequence alignments of Thermotogae MCPs; sequence alignments of CheA P5 domains; allele specific repressor mutations between Tsr and CheW mapped onto the upper MCP interaction site; contact sites among components of the ternary structure; MCP contacts to CheW and CheA determined by various means. This material is available free of charge via the Internet at <http://pubs.acs.org>.

Accession Codes

Deposited in the Protein Data Bank under accession code 4JPB.

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Notes

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■ ABBREVIATIONS USED

cryo-electron microscopy, cryoEM; FPLC, fast protein liquid chromatography; MCP, methyl-accepting chemotaxis protein; MCP_C, MCP cytoplasmic region; KCM, kinase control module; MR, molecular replacement; PDS, ESR, electron-spin resonance spectroscopy; P5, CheA regulator domain; PIR, protein interaction region; r, receptor; k, kinase; w, CheW; Tm14, *T. maritima* MCP TM0014; Tm14 PIR, Tm14s; TAM-IDS, tryptophan and alanine mutation to identify docking sites; Tris, 2-amino-2-hydroxymethyl-propane-1,3-diol

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